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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/813,483	03/29/2004	Jun Liu	P2026R1	5594	
9157	7590 12/09/2005		EXAM	EXAMINER	
GENENTECH, INC.			KIM, YU	KIM, YUNSOO	
1 DNA WAY SOUTH SAN FRANCISCO, CA 94080			ART UNIT	PAPER NUMBER	
	·		1644		

DATE MAILED: 12/09/2005

Please find below and/or attached an Office communication concerning this application or proceeding.

Advisory Action Before the Filing of an Appeal Brief

Application No.	Applicant(s)		
10/813,483	LIU ET AL.		
Examiner	Art Unit		
Yunsoo Kim	1644		

	T dribbo Tairi	1044	
The MAILING DATE of this communication app	ears on the cover sheet with the	correspondence ado	lress
THE REPLY FILED 07 November 2005 FAILS TO PLACE TH	IS APPLICATION IN CONDITION	FOR ALLOWANCE.	
1. The reply was filed after a final rejection, but prior to or of this application, applicant must timely file one of the folking places the application in condition for allowance; (2) a N a Request for Continued Examination (RCE) in compliant time periods:	owing replies: (1) an amendment, a otice of Appeal (with appeal fee) in the with 37 CFR 1.114. The reply	affidavit, or other evider n compliance with 37 C	nce, which FR 41.31; or (3)
a) The period for reply expiresmonths from the mailing			
b) The period for reply expires on: (1) the mailing date of this no event, however, will the statutory period for reply expire Examiner Note: If box 1 is checked, check either box (a) o	later than SIX MONTHS from the mai	ling date of the final reject	ion.
TWO MONTHS OF THE FINAL REJECTION. See MPEP		HE FINST KEPLT WAS F	TED WITHIN
Extensions of time may be obtained under 37 CFR 1.136(a). The dat have been filed is the date for purposes of determining the period of e under 37 CFR 1.17(a) is calculated from: (1) the expiration date of the set forth in (b) above, if checked. Any reply received by the Office lat may reduce any earned patent term adjustment. See 37 CFR 1.704(b) NOTICE OF APPEAL	e on which the petition under 37 CFR xtension and the corresponding amou shortened statutory period for reply or er than three months after the mailing	nt of the fee. The appropriginally set in the final Offi	iate extension fee ice action; or (2) as
2. The Notice of Appeal was filed on A brief in comfiling the Notice of Appeal (37 CFR 41.37(a)), or any ext	ension thereof (37 CFR 41.37(e)),	to avoid dismissal of th	hs of the date of ne appeal. Since
a Notice of Appeal has been filed, any reply must be file AMENDMENTS	a within the time period set forth ir	137 CFR 41.37(a).	
	but prior to the data of filing a bri	of will not be autored b	
 The proposed amendment(s) filed after a final rejection They raise new issues that would require further c 			ecause
(b) They raise the issue of new matter (see NOTE bel		OTE Delow),	
(c) They are not deemed to place the application in be		reducing or simplifying	the issues for
appeal; and/or	the form for appear by materially	readoning or simplifying	the issues for
(d) They present additional claims without canceling a	corresponding number of finally r	eiected claims.	
NOTE: (See 37 CFR 1.116 and 41.33(a))		•	
4. The amendments are not in compliance with 37 CFR 1.		Compliant Amendment	(PTOL-324)
5. Applicant's reply has overcome the following rejection(s			(*
6. Newly proposed or amended claim(s) would be a		a timely filed amendme	ant canceling the
non-allowable claim(s).	anovable ii dabiinted iii a deparat	o, amory mod amoramic	on canceling the
7. For purposes of appeal, the proposed amendment(s): all how the new or amended claims would be rejected is professed that the status of the claim(s) is (or will be) as follows:	will not be entered, or b) ☐ vovided below or appended.	will be entered and an e	explanation of
Claim(s) allowed:			
Claim(s) objected to: Claim(s) rejected: <u>1-17,20 and 22-27</u> .			
Claim(s) rejected. <u>1-17,20 and 22-27.</u> Claim(s) withdrawn from consideration: <u>18,19,21,28-45</u> and 18,19,21,28-45 and 18,19,21 and 18,1	and 48-50		
AFFIDAVIT OR OTHER EVIDENCE			
 The affidavit or other evidence filed after a final action, b because applicant failed to provide a showing of good awas not earlier presented. See 37 CFR 1.116(e). 	ut before or on the date of filing a nd sufficient reasons why the affid	Notice of Appeal will <u>no</u> avit or other evidence i	ot be entered s necessary and
9. The affidavit or other evidence filed after the date of filing entered because the affidavit or other evidence failed to showing a good and sufficient reasons why it is necessar	overcome all rejections under app	eal and/or appellant fa	ils to provide a
10. The affidavit or other evidence is entered. An explanation			
REQUEST FOR RECONSIDERATION/OTHER	and the states of the diaming after	and the second of all del	
 The request for reconsideration has been considered b <u>See Continuation Sheet.</u> 			nce because:
 Note the attached Information Disclosure Statement(s). 	(PTO/SB/08 or PTO-1449) Paper	No(s). <u>11/7/05</u>	
13. Other:			
		Patrick J. Nolan, Pl Primary Examiner Tech Center 1600,	
		· ·	

U.S. Patent and Trademark Office PTOL-303 (Rev. 7-05)

Continuation of 11. does NOT place the application in condition for allowance because:

Claims 1-17, 20, 22-27 stand rejected under 35 U.S.C. 103(a) as being unpatentable over Andya et al. (WO 97/04801, IDS ref. No.18, of record) in view of Relton et al. (WO 97/45140, of record), Kaisheva et al. (US2003/0113316, of record) and Merck Index (Merck Index, 10th Ed, 1983, p.797-798, of record) for the reasons set forth in the office action mailed 8/8/05.

Applicants' arguments filed on 11/7/05 have been fully considered but they are not persuasive.

Applicants argue that the Kaisheva reference is defective because it is not suitable for antibody concentration of 100-260mg/ml, limited to only combinations of cryprotectants, does not recognize increased turbidity by the addition of certain excipients and does not use glycine in the precise ratio of mannitol.

Contrary to Applicants' argument, Kaisheva reference is not limited to only the combinations of cryoprotectants. It is provided to show the well known use of arginine in the antibody formulation art. The stable antibody formulation of 100-260 mg/ml in histidine buffer was met by the Andya reference thus teachings of adding arginine at antibody formulation of greater than 50mg/ml further provides methods to improve antibody stability. Furthermore, applicant's comments regarding the required precise ratio of mannitol in glycine formulation is irrelevant as glycine in combination of mannitol is not claimed.

As is indicated in the previous office action mailed 8/8/05, the combination of the references of record teaches the claimed invention, the characterisitics of low turbidity and viscosity are expected property. Thus, it is examiner's position that the combinations of teachings (of record) remain obvious.

No claims are allowable.

PATRICK J. NOLAN, PH.D PRIMARY EXAMINER

farm John

12/7/05